Ref #	Hits	Search Query	DBs	Default Operator	Plurals	Time Stamp		
L1	15230	microbial and fermentation	US-PGPUB; USPAT; EPO; DERWENT	OR	ON	2005/08/31 20:17		
L2	420399	microbial and fermentation and recombiant "protein"	US-PGPUB; USPAT; EPO; DERWENT	OR	ON	2005/08/31 20:18		
L3	0	microbial and fermentation and "recombiant protein"	US-PGPUB; USPAT; EPO; DERWENT	OR	ON	2005/08/31 20:18		
L4	2467	microbial and fermentation and "recombinant protein"	US-PGPUB; USPAT; EPO; DERWENT	OR	ON	2005/08/31 20:32		
L5	37	microbial same fermentation same "recombinant protein"	US-PGPUB; USPAT; EPO; DERWENT	OR	OR ON 2005/0			
L6	1019	oscillatingly	US-PGPUB; USPAT; EPO; DERWENT	OR ·	ON	2005/08/31 20:19		
L7	35	oscillatingly same cycle	US-PGPUB; USPAT; EPO; DERWENT	OR	ON	2005/08/31 20:20		
L8	0	oscillatingly and fermentation	US-PGPUB; USPAT; EPO; DERWENT	OR	ON	2005/08/31 20:20		
L9	0	microbial and fermentation and "recombinant protein" and L6	US-PGPUB; OR USPAT; EPO; DERWENT		ON	2005/08/31 20:21		
L10	1971	microbial and fermentation and "recombinant protein" and period	US-PGPUB; USPAT; EPO; DERWENT	OR	ON	2005/08/31 20:21		
L11	1579	microbial and fermentation and "recombinant protein" and period and cycle	US-PGPUB; USPAT; EPO; DERWENT	OR	ON	2005/08/31 20:21		
L12	137	microbial and fermentation and "recombinant protein" and period same cycle	US-PGPUB; USPAT; EPO; DERWENT	OR	ON	2005/08/31 20:21		

L13	33	microbial same fermentation and "recombinant protein" and period same cycle	US-PGPUB; USPAT; EPO; DERWENT	OR	ON	2005/08/31 20:21
L14	1097	microbial and fermentation and "recombinant protein" and 435/69. 1.ccls.	US-PGPUB; USPAT; EPO; DERWENT	OR	ON	2005/08/31 20:32
L15	789	microbial and fermentation and "recombinant protein" and 530/350.ccls.	US-PGPUB; USPAT; EPO; DERWENT	OR	ON	2005/08/31 20:32

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=> microbial and fermentation and "recombinant protein"
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         1 FILE DRUGU
        12 FILE EMBASE
       116 FILE ESBIOBASE
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         1 FILE FROSTI
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         3 FILE IFIPAT
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        10 FILE JICST-EPLUS
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            FILE TOXCENTER
           FILE USPATFULL
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  67 FILES SEARCHED...
       157 FILE USPAT2
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          16 PASCAL
          14 CEABA-VTB
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F28	3	IFIPAT
F29	2	DISSABS
F30	2	WPIDS
F31	2	WPINDEX
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F33	1	CANCERLIT
F34	1	DDFU
F35	1	DRUGU
F36	1	FROSTI
F37	1	GENBANK
F38	1	PHIN

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=> microbial and fermentation and "recombinant protein" and period and cycle
L2 2 MICROBIAL AND FERMENTATION AND "RECOMBINANT PROTEIN" AND PERIOD
AND CYCLE

## => d ab bib

- L2 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2005 ACS on STN
- AB Combined transcriptome and proteome anal. was carried out to understand metabolic and physiol. changes of E. coli during high cell d. cultivation (HCDC). The expression of genes of TCA cycle enzymes, NADH dehydrogenase and ATPase, was up-regulated during the exponential fed-batch period and was down-regulated afterward. However, expression of most of the genes involved in glycolysis and pentose phosphate pathway was up-regulated at the stationary phase. The expression of most of amino acid biosynthesis genes was down-regulated as cell d. increased, which seems to be the major reason for the reduced specific productivity of recombinant proteins during

HCDC. The expression of chaperone genes increased with cell d., suggesting that the high cell d. condition itself can be stressful to the cells. Severe competition for O2 at high cell d. seemed to make cells use cytochrome bd, which is less efficient but has a higher O2 affinity than cytochrome bo3. Population cell d. itself strongly affected the expression of porin protein genes, especially ompF, and hence the permeability of the outer membrane. Expression of phosphate starvation genes was most strongly up-regulated toward the end of cultivation. It was also found that  $\sigma E$  (rpoE) plays a more important role than  $\sigma S$  (rpoS) at the stationary phase of HCDC. These findings should be invaluable in designing metabolic engineering and fermentation strategies for the production of recombinant proteins and metabolites by HCDC of E. coli.

- AN 2003:195554 CAPLUS
- DN 138:398487
- TI Combined transcriptome and proteome analysis of Escherichia coli during high cell density culture
- AU Yoon, Sung Ho; Han, Mee-Jung; Lee, Sang Yup; Jeong, Ki Jun; Yoo, Jong-Shin
- CS Metabolic and Biomolecular Engineering National Research Laboratory,
  Department of Chemical and Biomolecular Engineering, and BioProcess
  Engineering Research Center, Korea Advanced Institute of Science and
  Technology, Daejeon, 305-701, S. Korea
- SO Biotechnology and Bioengineering (2003), 81(7), 753-767 CODEN: BIBIAU; ISSN: 0006-3592
- PB John Wiley & Sons, Inc.
- DT Journal
- LA English
- RE.CNT 31 THERE ARE 31 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- => d ab bib

## NSWER 1 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN

- For many years, high broth viscosity has remained a key challenge in large-scale filamentous fungal fermns. In previous studies, we showed that broth viscosity could be reduced by pulsed addition of limiting carbon during fed-batch fermentation The objective in this study was to determine how changing the frequency of pulsed substrate addition affects fungal morphol., broth rheol., and recombinant enzyme productivity. To accomplish this, a series of duplicate fed-batch fermns. were performed in 20-L fermentors with a recombinant glucoamylase producing strain of Aspergillus oryzae. The total cycle time for substrate pulsing was varied over a wide range (30 - 2,700 s), with substrate added only during the first 30% of each cycle. As a control, a fermentation was conducted with continuous substrate feeding, and in all fermns. the same total amount of substrate was added. Results show that the total biomass concentration remained relatively unaltered, while a substantial decrease in the mean projected area of fungal elements (i.e., average size) was observed with increasing cycle time. This led to reduced broth viscosity and increased oxygen uptake rate. However, high values of cycle time (i.e., 900 - 2,700 s) showed a significant increase in fungal conidia formation and significantly reduced recombinant enzyme productivity, suggesting that the fungi channeled substrate to storage compds. rather than to recombinant protein. In addition to explaining the effect of cycle time on fermentation performance, these results may aid in explaining the discrepancies observed on scale-up to larger fermentors.
- AN 2005:195802 CAPLUS
- DN 142:446127
- TI Effect of **cycle** time on fungal morphology, broth rheology, and recombinant enzyme productivity during pulsed addition of limiting carbon source
- AU Bhargava, Swapnil; Wenger, Kevin S.; Rane, Kishore; Rising, Vanessa; Marten, Mark R.
- CS Department of Chemical and Biochemical Engineering, University of Maryland, Baltimore County, Baltimore, MD, 21250, USA
- SO Biotechnology and Bioengineering (2005), 89(5), 524-529 CODEN: BIBIAU; ISSN: 0006-3592
- PB John Wiley & Sons, Inc.
- DT Journal
- LA English
- RE.CNT 47 THERE ARE 47 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

## => d ti 1-47

- L3 ANSWER 1 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Effect of cycle time on fungal morphology, broth rheology, and recombinant enzyme productivity during pulsed addition of limiting carbon source
- L3 ANSWER 2 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Flocculation gene expression in genetically modified microorganisms is regulated by medium changes and aids separation of microbes and fermentation products
- L3 ANSWER 3 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Combined transcriptome and proteome analysis of Escherichia coli during high cell density culture
- L3 ANSWER 4 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Limiting factors in Escherichia coli fed-batch production of recombinant proteins

- L3 ANSWER 5 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Metabolic adaptation of Escherichia coli during temperature-induced recombinant protein production: 2. Redirection of metabolic fluxes
- L3 ANSWER 6 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN
- TI A comparative study of global stress gene regulation in response to overexpression of **recombinant proteins** in Escherichia coli
- L3 ANSWER 7 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN
- TI 13C NMR Evidence for Pyruvate Kinase Flux Attenuation Underlying Suppressed Acid Formation in Bacillus subtilis
- L3 ANSWER 8 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Metabolic flux distributions in recombinant Saccharomyces cerevisiae during foreign protein production
- L3 ANSWER 9 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Metabolic pathway analysis of recombinant Saccharomyces cerevisiae with a galactose-inducible promoter based on a signal flow modeling approach
- L3 ANSWER 10 OF 12 MEDLINE on STN
- TI Influence of controlled glucose oscillations on a fed-batch process of recombinant Escherichia coli.
- L3 ANSWER 11 OF 12 MEDLINE on STN
- TI Use of a cell recycle reactor to increase production of a proteolysis-susceptible peptide secreted from recombinant Saccharomyces cerevisiae.
- L3 ANSWER 12 OF 12 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on
- TI Limiting factors in Escherichia coli fed-batch production of recombinant proteins.
- => d ab bib 1-12
- L3 ANSWER 1 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN
- For many years, high broth viscosity has remained a key challenge in AB large-scale filamentous fungal fermns. In previous studies, we showed that broth viscosity could be reduced by pulsed addition of limiting carbon during fed-batch fermentation The objective in this study was to determine how changing the frequency of pulsed substrate addition affects fungal morphol., broth rheol., and recombinant enzyme productivity. To accomplish this, a series of duplicate fed-batch fermns. were performed in 20-L fermentors with a recombinant glucoamylase producing strain of Aspergillus oryzae. The total cycle time for substrate pulsing was varied over a wide range (30 - 2,700 s), with substrate added only during the first 30% of each cycle. As a control, a fermentation was conducted with continuous substrate feeding, and in all fermns. the same total amount of substrate was added. Results show that the total biomass concentration remained relatively unaltered, while a substantial decrease in the mean projected area of fungal elements (i.e., average size) was observed with increasing cycle time. This led to reduced broth viscosity and increased oxygen uptake rate. However, high values of cycle time (i.e., 900 - 2,700 s) showed a significant increase in fungal conidia formation and significantly reduced recombinant enzyme productivity, suggesting that the fungi channeled substrate to storage compds. rather than to recombinant protein. In addition to explaining the effect

of cycle time on fermentation performance, these results may aid in explaining the discrepancies observed on scale-up to larger fermentors.

AN 2005:195802 CAPLUS

DN 142:446127

- TI Effect of cycle time on fungal morphology, broth rheology, and recombinant enzyme productivity during pulsed addition of limiting carbon source
- AU Bhargava, Swapnil; Wenger, Kevin S.; Rane, Kishore; Rising, Vanessa; Marten, Mark R.
- CS Department of Chemical and Biochemical Engineering, University of Maryland, Baltimore County, Baltimore, MD, 21250, USA
- SO Biotechnology and Bioengineering (2005), 89(5), 524-529 CODEN: BIBIAU; ISSN: 0006-3592
- PB John Wiley & Sons, Inc.
- DT Journal
- LA English
- RE.CNT 47 THERE ARE 47 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L3 ANSWER 2 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN
- This invention relates to flocculation gene expression in genetically AΒ modified microorganisms, regulated by medium changes, that aids separation of microbes and fermentation products. While the invention focuses on modified yeast strains (hundreds of Saccharomyces cerevisiae strains are claimed), modification of bacteria, fungus, archae, alga and protozoa are also included. The target of genetic engineering in this invention is a series of flocculation genes, including genes FLO2-FLO11, Lg-FLO1, FLO1S, FLOIL, PKC1, sfllou, fsulou, fsulou, tuplou, cyc8ou, cka2, and FMC1. Regulation of the genes is provided by gene promoters responsive to environmental cues, specifically promoters from genes Mox, HSP30p, pMET3, and ADH. The triggers for regulation of gene expression are provided in the cell culture media at the end of a fermentation cycle, including fall of pH, thermal shock, change in sugar, nitrogen or ethanol composition, or phys. excitation. Upon stimulation of the environmental cue, flocculation gene expression is modified, causing the cells to settle quickly to the bottom of the fermentation vat, facilitating separation of microbes and fermentation product. This invention can be applied to production of a wide range of food or pharmaceutical products including alc. beverages and solution of recombinant proteins.
- AN 2004:41622 CAPLUS
- DN 140:106559
- TI Flocculation gene expression in genetically modified microorganisms is regulated by medium changes and aids separation of microbes and fermentation products
- IN Gomes de Souza, Marcos; Pereira Junior, Haroldo Alves
- PA Salinbar S.A., Urug.
- SO PCT Int. Appl., 77 pp.

CODEN: PIXXD2

- DT Patent
- LA English
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			GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	ΚZ,	LC,	LK,	LR,
			LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NI,	NO,	ΝZ,	OM,
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FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG BR 2002003754 20040525 BR 2002-3754 20020708 Α EP 1551951 A1 20050713 EP 2003-735214 20030708 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, R: IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK PRAI BR 2002-3754 20020708 А WO 2003-BR89 W 20030708 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 4 ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 3 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN

AB Combined transcriptome and proteome anal. was carried out to understand metabolic and physiol. changes of E. coli during high cell d. cultivation (HCDC). The expression of genes of TCA cycle enzymes, NADH dehydrogenase and ATPase, was up-regulated during the exponential fed-batch period and was down-regulated afterward. However, expression of most of the genes involved in glycolysis and pentose phosphate pathway was up-regulated at the stationary phase. The expression of most of amino acid biosynthesis genes was down-regulated as cell d. increased, which seems to be the major reason for the reduced specific productivity of recombinant proteins during HCDC. The expression of chaperone genes increased with cell d., suggesting that the high cell d. condition itself can be stressful to the cells. Severe competition for 02 at high cell d. seemed to make cells use cytochrome bd, which is less efficient but has a higher O2 affinity than cytochrome bo3. Population cell d. itself strongly affected the expression of porin protein genes, especially ompF, and hence the permeability of the outer membrane. Expression of phosphate starvation genes was most strongly up-regulated toward the end of cultivation. It was also found that  $\sigma E$  (rpoE) plays a more important role than  $\sigma S$  (rpoS) at the stationary phase of HCDC. These findings should be invaluable in designing metabolic engineering and fermentation strategies for the production of recombinant proteins and metabolites by HCDC of E. coli.

AN 2003:195554 CAPLUS

DN 138:398487

TI Combined transcriptome and proteome analysis of Escherichia coli during high cell density culture

AU Yoon, Sung Ho; Han, Mee-Jung; Lee, Sang Yup; Jeong, Ki Jun; Yoo, Jong-Shin

CS Metabolic and Biomolecular Engineering National Research Laboratory, Department of Chemical and Biomolecular Engineering, and BioProcess Engineering Research Center, Korea Advanced Institute of Science and Technology, Daejeon, 305-701, S. Korea

SO Biotechnology and Bioengineering (2003), 81(7), 753-767 CODEN: BIBIAU; ISSN: 0006-3592

PB John Wiley & Sons, Inc.

DT Journal

LA English

RE.CNT 31 THERE ARE 31 CITED REFERENCES AVAILABLE FOR THIS RECORD

- L3 ANSWER 4 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN
- Fed-batch production of recombinant  $\beta$ -galactosidase in E. coli was AB studied with respect to the specific growth rate at induction. The cultivations were designed to induce protein production by IPTG at a glucose feed rate corresponding to high  $(\mu = 0.5 \text{ h-1})$  or low  $(\mu = 0.1 \text{ h-1})$ specific growth rate. Protein production rate was approx. 100% higher at the higher specific growth rate, resulting in the accumulation of  $\beta$ -galactosidase up to 30% of the total cell protein. Transcription anal. showed that  $\beta$ -galactosidase-specific mRNA was immediately formed after induction (<5 min), but the amount was the same in both cases and was thus not the initial limiting factor. The content of ribosomes, as represented by rRNA, rapidly decreased with specific growth rate from a relative level of 100%, at the high specific growth rate, to 20% at the low specific growth rate. At high specific growth rate, ribosomes were addnl. degraded upon induction due to the high production level. Translation therefore seemed to be the initial limiting factor of the protein synthesis capacity. The alarmone, guanosine tetraphosphate increased at both high and low feed level inductions, indicating an induction-forced starvation of charged tRNA and/or glucose. The altered physiol. status was also detected by the formation of acetic acid. However, the higher production rate resulted in high-level accumulation of acetic acid, which was absent at low feed rate production Acetic acid production is thus coupled to

the

high product formation rate and is proposed to be due either to a precursor drain of Krebs cycle intermediates and a time lag before induction of the glyoxalate shunt, or to single amino acid overflow, since the model product is relatively poor in glycine and alanine. In conclusion, it is proposed that production at high specific growth rate becomes precursor-limited, while production at low specific growth rate is carbon- and/or energy-limited.

- AN 2003:12171 CAPLUS
- DN 138:220444
- TI Limiting factors in Escherichia coli fed-batch production of recombinant proteins
- AU Sanden, Anna Maria; Prytz, Ingela; Tubulekas, Ioannis; Forberg, Cecilia; Le, Ha; Hektor, Andrea; Neubauer, Peter; Pragai, Zoltan; Harwood, Colin; Ward, Alan; Picon, Antonia; Teixeira de Mattos, Joost; Postma, Pieter; Farewell, Anne; Nystrom, Thomas; Reeh, Solvejg; Pedersen, Steen; Larsson, Gen
- CS The Swedish Centre for Bioprocess Technology, Stockholm Center for Physics, Astronomy and Biotechnology, Stockholm, SE-106 91, Swed.
- SO Biotechnology and Bioengineering (2003), 81(2), 158-166 CODEN: BIBIAU; ISSN: 0006-3592
- PB John Wiley & Sons, Inc.
- DT Journal
- LA English
- RE.CNT 27 THERE ARE 27 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L3 ANSWER 5 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN
- AB The impact of temperature-induced synthesis of human basic fibroblast growth factor (hFGF-2) in high-cell-d. cultures of recombinant Escherichia coli was studied by estimating metabolic flux variations. Metabolic flux distributions in E. coli were calculated by means of a stoichiometric network and linear programming. After the temperature upshift, a substantially elevated

energy demand for synthesis of hFGF-2 and heat shock proteins resulted in a redirection of metabolic fluxes. Catabolic pathways like the Embden-Meyerhof-Parnas pathway and the tricarboxylic acid (TCA) cycle showed significantly enhanced activities, leading to reduced flux to growth-associated pathways like the pentose phosphate pathway and other anabolic pathways. Upon temperature upshift, an excess of NADPH was

produced in the TCA **cycle** by isocitrate dehydrogenase. The metabolic model predicted the involvement of a transhydrogenase generating addnl. NADH from NADPH, thereby increasing ATP regeneration in the respiratory chain. The influence of the temperature upshift on the host's metabolism was investigated by means of a control strain harboring the "empty" parental expression vector. The metabolic fluxes after the temperature upshift were redirected similarly to the production strain; the effects, however, were observed to a lesser extent and with different time profiles.

AN 2002:796103 CAPLUS

DN 138:23758

- TI Metabolic adaptation of Escherichia coli during temperature-induced recombinant protein production: 2. Redirection of metabolic fluxes
- AU Weber, Jan; Hoffmann, Frank; Rinas, Ursula
- CS Biochemical Engineering Division, GBF German Research Center for Biotechnology, Braunschweig, 38124, Germany
- SO Biotechnology and Bioengineering (2002), 80(3), 320-330 CODEN: BIBIAU; ISSN: 0006-3592
- PB John Wiley & Sons, Inc.
- DT Journal
- LA English
- RE.CNT 39 THERE ARE 39 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L3 ANSWER 6 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN
- Global gene regulation throughout the Escherichia coli stress response to AΒ overexpression of each of five recombinant proteins was evaluated. Reverse-transcriptase polymerase chain reaction-amplified mRNA from induced and control cells were hybridized with a DNA array of Kohara clones representing 16% (700 genes) of the E. coli genome. Subsequently, Northern anal. was performed for quantification of specific gene dynamics and statistically significant overlap in the regulation of 11 stress-related genes was found using correlation anal. The results reported here establish that there are dramatic changes in the transcription rates of a broad range of stress genes (representing multiple regulons) after induction of recombinant protein. Specifically, the responses included significantly increased upregulation of heat shock (ftsH, clpP, lon, ompT, degP, groEL, aceA, ibpA), SOS/DNA damage (recA, lon, IS5 transposase), stationary phase (rpoS, aceA), and bacteriophage life cycle (ftsH, recA) genes. Importantly, similarities at the microscopic (gene) level were not clearly reflected at the macroscopic (growth rate, lysis) level. The use of such dynamic data is critical to the design of gene-based sensors, the engineering of metabolic pathways, and the determination of parameters (harvest and induction

times) needed for successful recombinant E. coli fermns. (c) 2000 Academic Press.

- AN 2000:903910 CAPLUS
- DN 135:191149
- TI A comparative study of global stress gene regulation in response to overexpression of recombinant proteins in Escherichia coli
- AU Gill, R. T.; Valdes, J. J.; Bentley, W. E.
- CS Department of Chemical Engineering, University of Maryland, College Park, MD, 20742, USA
- SO Metabolic Engineering (2000), 2(3), 178-189 CODEN: MEENFM; ISSN: 1096-7176
- PB Academic Press
- DT Journal
- LA English
- RE.CNT 54 THERE ARE 54 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L3 ANSWER 7 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN
- When batch and continuous Bacillus subtilis cultures are provided with a AB small amount of citrate, acid production ceases, carbon yield increases by more than 2-fold, and the productivity of recombinant protein increases. It has been hypothesized that pyruvate kinase activity is attenuated, which in turn lowers glucose flux and minimizes the acid overflow prompted by low Krebs cycle capacity. To complement existing enzyme activity, linear programming, and metabolite pool studies, 13C NMR studies were performed. Atom mapping and isotopomer mapping matrix methods were used to select the best glucose label. "Best" was defined such that the NMR spectra of glutamate associated with metabolizing labeled glucose via the different candidate metabolic trafficking scenarios would differ considerably in fine structure (e.g., relative singlet intensities). When expts. were performed with [1-13C]glucose, the observed NMR spectra corresponded well to the one predicted to arise when the metabolic trafficking occurs according to a pyruvate kinase attenuation scenario. This evidence further fortifies the prospects for successfully basing a metabolic engineering strategy on reducing pyruvate kinase activity to better match glycolytic and Krebs cycle capacities.
- AN 2000:186222 CAPLUS
- DN 132:307338
- TI 13C NMR Evidence for Pyruvate Kinase Flux Attenuation Underlying Suppressed Acid Formation in Bacillus subtilis
- AU Phalakornkule, C.; Fry, B.; Zhu, T.; Kopesel, R.; Ataai, M. M.; Domach, M. M.
- CS Department of Chemical Engineering Biotechnology & Health Engineering Program, Carnegie Mellon University, Pittsburgh, PA, USA
- SO Biotechnology Progress (2000), 16(2), 169-175 CODEN: BIPRET; ISSN: 8756-7938
- PB American Chemical Society
- DT Journal
- LA English
- RE.CNT 13 THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L3 ANSWER 8 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN
- A stoichiometric flux balancing anal. was applied to the recombinant yeast AB cultivation to examine the cellular physiol. and relationship between the production of heterologous protein and metabolic fluxes. The fluxes in the metabolic pathway within a recombinant S. cerevisiae grown on galactose alone or mixts. of galactose and ethanol medium were calculated It is found that an amplification of the PP (pentose phosphate) pathway activity resulted in an improvement of the foreign protein expression and cell yield on ATP. The carbon source used for biosynthesis from TCA cycle in the exponential growth phase was 2 and 5-fold higher, resp., as compared with that in the late exponential growth phase and stationary phase in batch culture with galactose min. medium. The metabolism of ethanol together with galactose in the recombinant cell looks like increasing the flux from Acetyl-CoA to TCA cycle, and amplifying the flux directing the synthesis of various kinds of precursors such as amino acids and nucleic acid which are necessary for production of a foreign protein. Metabolic flux distribution anal. also shows that the ATP synthesis rate under substrate-level phosphorylation in the mixed carbon source cultivation was lower than that in the sole carbon source (galactose) during the expression of foreign protein. However, the total ATP production rate was higher in the mixed carbon source cultivation.
- AN 1997:441121 CAPLUS
- DN 127:148372
- TI Metabolic flux distributions in recombinant Saccharomyces cerevisiae during foreign protein production
- AU Jin, Sha; Ye, Kaiming; Shimizu, Kazuyuki
- CS Department of Biochemical Engineering and Science, Kyushu Inst. Technol., Fukuoka, 820, Japan

- Journal of Biotechnology (1997), 54(3), 161-174 CODEN: JBITD4; ISSN: 0168-1656
- PB Elsevier
- DT Journal
- English LΑ
- THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 24 ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L3 ANSWER 9 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN
- AB The objective of this work was to develop a signal flow diagram-based modeling approach proposed by Endo et al. (1976) to organize the network of complex metabolic reactions occurring in the living cell, employing directed signal flow diagram in which the enzyme reaction coefficient was defined as the metabolic transfer coefficient Using this concept, a metabolic reaction between substrate A and product B could be regarded as a signal transmittance from A to B. A very simple set of linear equations was then derived to analyze the flow directions of the carbon fluxes and the degree of activation of certain metabolic pathways within a recombinant yeast, in which the specific consumption rate of galactose, the specific rate of change of ethanol and the specific uptake rate of oxygen constituted the three input nodes of the metabolic signal transfer system. The output nodes were the specific growth rate of the cells, the specific production rate of the recombinant protein and the specific evolution rate of carbon dioxide. In this way, the effect of the culture conditions on cell growth and recombinant protein production under control of the GAL10 promoter could be characterized in terms of the metabolic pathways based on observable variables such as the cell and product concns., and the carbon dioxide and oxygen in the exhaust gas. This approach was successfully applied to an anal. of the metabolic pathways occurring in the fed-batch cultivation of a recombinant yeast where galactose served as both the carbon source for cell growth and as an inducer for expression of the recombinant gene. The expression period was classified into three phases - named the switch, expression, and stationary phases - on the basis of calcns. using signal flow equations. In the switch phase, biomaterials for cell growth were found to be synthesized through the TCA cycle and UR loop, and less galactose entered the induction pathway. In the stationary phase, on the other hand, the formation of biomaterials for cell growth occurred mainly through the PP and EMP pathways and the TCA cycle. The degree of activation of the induction pathway was reduced in this phase. in either of the previous two phases, in the expression phase a significant amount of galactose was directed towards the induction pathway to stimulate the expression of the recombinant gene, and the PP pathway played the major role in the synthesis of biomaterials for cell growth. ΑN
- 1996:34349 CAPLUS
- 124:115511 DN
- Metabolic pathway analysis of recombinant Saccharomyces cerevisiae with a ΤI galactose-inducible promoter based on a signal flow modeling approach
- ΑU Jin, Sha; Ye, Kaiming; Shimizu, Kazuyuki
- Dep. of Biochemical Engineering and Science, Kyushu Inst. of Technology, CS Fukuoka, 820, Japan
- Journal of Fermentation and Bioengineering (1995), 80(6), 541-51 SO CODEN: JFBIEX; ISSN: 0922-338X
- PB Society for Fermentation and Bioengineering, Japan
- DT Journal
- LΑ English
- L3 ANSWER 10 OF 12 MEDLINE on STN
- The influence of glucose oscillations on cell growth and product formation AΒ of a recombinant Escherichia coli culture producing a heterologous alpha-glucosidase was studied in fed-batch cultures in a laboratory bioreactor. Glucose oscillations were created by an on/off-feeding mode in either fast cycles (1 min) or slow cycles (4 min)

and compared to a process with constant glucose addition. The study indicates that glucose oscillations influence the product stability and the overgrowth of plasmid-free cells if such cultures are not performed under continuous pressure for selection of plasmid-containing cells. Although the glucose uptake capacity decreased after induction of the recombinant alpha-glucosidase in all cultures performed, the up-growth of plasmid-free cells during the production phase was strongly inhibited by fast oscillations. In contrast, plasmid-free cells grew up when constant feeding or slow cycles were applied. Our data suggest that the various feed protocols effect the specific carbon dioxide formation rate differently, with the highest production of carbon dioxide in the cultivations with fast cycles. In connection to product formation the initial alpha-glucosidase accumulation was the same in all cultures, but the stability of the product was significantly lower in the cultivation with slow cycles. Our results from laboratory experiments are discussed in relation to the mixing situation in large-scale bioreactors.

- AN 2000275040 MEDLINE
- DN PubMed ID: 10817339
- TI Influence of controlled glucose oscillations on a fed-batch process of recombinant Escherichia coli.
- AU Lin H Y; Neubauer P
- CS Martin-Luther-Universitat Halle-Wittenberg, Fachbereich Biochemie /Biotechnologie, Institut fur Biotechnologie, Halle, Germany.
- SO Journal of biotechnology, (2000 Apr 14) 79 (1) 27-37. Journal code: 8411927. ISSN: 0168-1656.
- CY Netherlands
- DT Journal; Article; (JOURNAL ARTICLE)
- LA English
- FS Priority Journals
- EM 200007
- ED Entered STN: 20000714

Last Updated on STN: 20000714 Entered Medline: 20000706

- L3 ANSWER 11 OF 12 MEDLINE on STN
- AB Operation of a continuous microbial fermentor with cell recycle can significantly reduce degradation-associated loss of a secreted protein product. Under continuous fermentation conditions, proteolysis of a recombinant growth hormone releasing factor (GRF) analog secreted by S. cerevisiae was first order with respect to GRF concentration. The maximal GRF concentration was increased from 5 mg/l to 30 mg/l by the use of a cell recycle reactor, and volumetric productivity was increased more than 10-fold to an average of 10 mg/l-1/h-1. A mathematical model shows that increased productivity in the cell recycle reactor results from a reduced degradation rate and a shorter residence time of the product in the fermentor.
- AN 91025893 MEDLINE
- DN PubMed ID: 1366627
- TI Use of a cell recycle reactor to increase production of a proteolysis-susceptible peptide secreted from recombinant Saccharomyces cerevisiae.
- AU Siegel R S; Brierley R A
- CS Salk Institute Biotechnology/Industrial Associates, Inc. (SIBIA), San Diego, CA 92138.
- SO Bio/technology (Nature Publishing Company), (1990 Jul) 8 (7) 639-43. Journal code: 8309273. ISSN: 0733-222X.
- CY United States
- DT Journal; Article; (JOURNAL ARTICLE)
- -LA English
- FS Biotechnology
- EM 199012
- ED Entered STN: 19950809

Last Updated on STN: 19950809 Entered Medline: 19901214

- L3 ANSWER 12 OF 12 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on STN
- Fed-batch production of recombinant beta-galactosidase in E. coli was AB studied with respect to the specific growth rate at induction. The cultivations were designed to induce protein production by IPTG at a glucose feed rate corresponding to high (mu=0.5 h-1) or low (mu=0.1 h-1) specific growth rate. Protein production rate was approximately 100% higher at the higher specific growth rate, resulting in the accumulation of beta-galactosidase up to 30% of the total cell protein. Transcription analysis showed that beta-galactosidase-specific messenger RNA was immediately formed after induction (<5 min), but the amount was the same in both cases and was thus not the initial limiting factor. The content of ribosomes, as represented by rRNA, rapidly decreased with specific growth rate from a relative level of 100%, at the high specific growth rate, to 20% at the low specific growth rate. At high specific growth rate, ribosomes were additionally degraded upon induction due to the high production level. Translation therefore seemed to be the initial limiting factor of the protein synthesis capacity. The alarmone guanosine tetraphosphate increased at both high and low feed level inductions, indicating an induction-forced starvation of charged tRNA and/or glucose. The altered physiological status was also detected by the formation of acetic acid. However, the higher production rate resulted in high-level accumulation of acetic acid, which was absent at low feed rate production. Acetic acid production is thus coupled to the high product formation rate and is proposed to be due either to a precursor drain of Krebs cycle intermediates and a time lag before induction of the glyoxalate shunt, or to single amino acid overflow, since the model product is relatively poor in glycin and alanin. In conclusion, it is proposed that production at high specific growth rate becomes precursor-limited, while production at low specific growth rate is carbonand/or energy-limited.
- AN 2003:73731 BIOSIS
- DN PREV200300073731
- TI Limiting factors in Escherichia coli fed-batch production of recombinant proteins.
- AU Sanden, Anna Maria; Prytz, Ingela; Tubulekas, Ioannis; Forberg, Cecilia; Le, Ha; Hektor, Andrea; Neubauer, Peter; Pragai, Zoltan; Harwood, Colin; Ward, Alan; Picon, Antonia; de Mattos, Joost Teixeira; Postma, Pieter; Farewell, Anne; Nystrom, Thomas; Reeh, Solvejg; Pedersen, Steen; Larsson, Gen [Reprint Author]
- CS Stockholm Center for Physics, Astronomy and Biotechnology, Swedish Centre for Bioprocess Technology, SE-106 91, Stockholm, Sweden Gen.Larsson@biotech.kth.se
- SO Biotechnology and Bioengineering, (January 20 2003) Vol. 81, No. 2, pp. 158-166. print. CODEN: BIBIAU. ISSN: 0006-3592.
- DT Article
- LA English
- ED Entered STN: 29 Jan 2003 Last Updated on STN: 29 Jan 2003

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